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Lab-on-a-disc platform for screening of genetically modified E. coli cells via cell-free electrochemical detection of \( p \)-Coumaric acid

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Abstract

We present a robust easy to use lab-on-a-disc (LoD) device with integrated sample pre-treatment and electrochemical detection system for cell-free detection of a secondary metabolite, \( p \)-Coumaric acid (\( pHCA \)), produced by genetically modified \( E. \) coli. In the LoD device, which incorporates eight filtration and electrochemical detection units, the sample filtration was performed by rotating the disc using a programmable closed-loop stepper motor. The electrodes, patterned on plastic substrate, were connected through a printed circuit board to the slip ring using a robust magnetic clamping system that enables easy assembly and robust electrical connections. \( pHCA \) was quantified in a linear range from 0.125 up to 2 mM using square wave voltammetry. The platform was successfully used for the quantification of \( pHCA \) produced by two genetically modified \( E. \) coli strains after 24 h of cell culture. The data obtained from the electrochemical measurements showed good correlation with high performance liquid chromatographic analysis. The developed LoD system offers fast and easy detection of \( pHCA \), enabling screening of genetically modified organisms based on the quantity of produced secondary metabolites.

Keywords: lab-on-a-disc; cell free detection; electrochemical sensor; bacterial metabolite; magnetic clamping; electrode on plastic substrate
1. Introduction

In the past few years, the interest for plant secondary metabolites has increased significantly due to their diverse biological functions and wide range of applications in chemical, food, health, cosmetic, and pharmaceutical industries (Balasundram et al., 2006; Barbulova et al., 2015; Soto et al., 2015; Sytar Oksana, 2012). \( p \)-Coumaric acid (\( pHCA \)) is one of these secondary metabolites with many physiological effects, acting as, e.g., a potent antioxidant, antimicrobial, anxiolytic, analgesic, antimutagen, sedative and immunoregulatory agent (Boz, 2015; Cheng et al., 2007; Ou et al., 2012). Unfortunately, extraction from plants requires a significant amount of starting material, harsh conditions and results in a limited purity of products (Ou et al., 2012). To overcome these challenges, scientists have altered the genome of microorganisms to develop metabolically engineered microbial strains for a more economical production of \( pHCA \) from inexpensive amino acids (Calero et al., 2016; Rodriguez et al., 2015; Xue et al., 2014). Biosynthesis of \( pHCA \) can be carried out via deamination of tyrosine (Tyr) by incorporating tyrosine ammonia lyase (TAL) gene in \( E.\ coli \) genome (Jendresen et al., 2015). Efficient optimization of the tyrosine pathway as well as the activity of the TAL enzyme for production of \( pHCA \) is generally validated by the screening of the genetically modified strains by quantifying the amount of produced \( pHCA \). High performance liquid chromatography (HPLC), gas chromatography, thin-layer chromatography and spectrophotometric assays are commonly used to quantify secondary metabolites from plants (Khoddami et al., 2013). However, there is a great need for novel platforms that allow high-throughput analysis with minimal sample handling for cost-effective, easy and fast screening of modified strains.

Centrifugal microfluidic devices, often referred to as lab-on-a-disc (LoD) systems, have gained interest in both academic research and industry (Burger et al., 2016; Smith et al., 2016). In a LoD platform, the centrifugal forces, induced by rotating the device, enable the transport and manipulation of liquids; therefore offer an alternative to traditional pressure-driven microfluidic systems. They require minimal instrumentation for liquid transport and handling, consequently enabling the development of a simple, compact and integrated detection systems. Although optical detection is the most commonly implemented approach on LoD devices (Donolato et al., 2015; Grumann et al., 2006; Kim et al., 2013; Koh et al., 2015; Uddin et al., 2016; Zhang et al., 2015), there are examples describing the integration of electrochemical (Abi-Samra et al., 2013; Amato et al., 2015; Andreasen et al., 2015; Cho et al., 1982; Kim et al., 2013; Li et al., 2013; Nwankire et al., 2015; Rattanarat et al., 2015) and mechanical (Bache et al., 2013; Bosco et al., 2011) sensors. Integration of electrochemical detection with microfluidic devices offers advantages, such as possibility to miniaturize both the transducer (electrode) and instrumentation as well as multiplexing and automation of measurements without losing the performance (Nyholm, 2005; Sassa et al., 2008; Xu et al., 2009). In addition, electrochemical detection is affected by neither sample turbidity nor optical path length, and it is possible to fabricate low-cost detection systems with minimal power demand (Felipe et al., 2014; Xu et al., 2011). However, direct detection of analytes in a complex sample matrix can be challenging when the potential interfering compounds and the possibility for surface fouling of the transducer are taken into
consideration. There are several reports on the integration of various sample handling (e.g. mixing, dilution) and pre-treatment methods (e.g. DNA extraction, blood separation, cell extraction from plasma) for LoD based applications (Grumann et al., 2006; Kim et al., 2014; Li et al., 2013; Nwankire et al., 2015) to improve detection accuracy.

In this work, we have developed a simple, robust and easy to use LoD platform with eight electrode arrays fabricated on a plastic substrate to facilitate electrochemical detection of pHCA in supernatants derived from E. coli cultures. To provide cell-free samples to the electrochemical detection units the LoD device had an integrated microporous filter membrane for simple sample pre-treatment. The platform was used for quantification of pHCA produced by two genetically modified E. coli strains.

2. Material and methods

2.1 Chemicals

Potassium hexacyanoferrate(II) (ferrocyanide), potassium hexacyanoferrate(III) (ferricyanide), potassium hydroxide (KOH), phosphate buffered saline (PBS), pHCA, the components of the culture medium for E. coli (minimal M9 medium) and the analytes used for the HPLC analysis were purchased from Sigma-Aldrich® Co. (St. Louis, MO, USA) unless otherwise stated. Minimal M9 medium, containing 6.8 g/L Na2HPO4·7H2O, 3.0 g/L KH2PO4, 0.5 g/L NaCl, 1.0 g/L NH4Cl, 2 mM MgSO4, 0.1 mM CaCl2, and 1:2000 (v/v) dilution of a trace element stock solution (10 g/L FeCl3 ·6H2O, 2 g/L ZnSO4·7H2O, 0.4 g/L CuCl2·2H2O, 1 g/L MnSO4·H2O, 0.6 g/L CoCl2·6H2O, and 3.2 mL/L of 0.5 M EDTA at pH 8), was prepared in water and autoclaved. 10 g/L glucose, 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG), and Wolfe’s Vitamin solution (purchased from ATCC® (LGC Standards, UK) and containing 10 μg/L pyridoxine-HCl, 5 μg/L thiamine-HCl, 5 μg/L riboflavin, 5 μg/L nicotinic acid, 5 μg/L calcium D-(+)-pantothenate, 5 μg/L p-aminobenzoic acid, 2 μg/L biotin, 2 μg/L folic acid, 5 μg/L lipoic acid, and 0.1 μg/L vitamin B12) were added to the autoclaved medium. In addition 2 mM Tyr was added as a substrate for pHCA production during cell culture. To maintain the plasmids in the developed E. coli strains, antibiotics, chloramphenicol and spectinomycin, were also added to the M9 medium at the final concentration of 34 mg/L and 50 mg/L, respectively and. M9 buffer was prepared using 6.8 g/L Na2HPO4·7H2O, 3.0 g/L KH2PO4, and 0.5 g/L NaCl. Ethanol (EtOH) 99% was purchased from VWR Chemicals (Søborg, Denmark). All the aqueous solutions were prepared in ultrapure water obtained from a Milli-Q® water purification system (Millipore Corporation, Billerica, MA, USA).

2.2 E. coli culture

E. coli control (CBJ786) and pHCA producing strains (CBJ792 and CBJ800) are derived as previously described (Jendresen et al., 2015) from the expression strain BL21(DE3)pLysS (Invitrogen/Life Technologies), carrying either the extrachromosomal plasmid pCDFDuet-1 (CBJ786) or derivative plasmids encoding for one of the tyrosine ammonia-lyases R_XAL (CBJ792) or FjTAL (CBJ800). For pHCA production assays, the E. coli strains were grown in 180 mL of M9 medium in 500 mL-flasks in
a temperature-controlled orbital shaker (30 °C, 250 rpm). 30 mL samples were withdrawn at 24 h, and supernatants were obtained by centrifugation (10 min at 10000 g, 4 °C). The optical density (OD) at 600 nm was measured in cuvettes with 1 cm path length using a UV-1600PC spectrophotometer (VWR, Radnor, Pennsylvania, USA).

2.3 Experimental setup
The fully assembled experimental set-up shown in Figure 1A is comprised of the LoD device mounted on the spin-stand, which consists of an electronic triggering system for the high-resolution camera (pco.pixelfly, PCO AG, Kelheim, Germany) and a spinning motor (Maxon motor, model RE 35) controlled by custom-made software in a LabVIEW environment. The electrodes (Figure 2B) incorporated in the fluidic platform were connected to a CHI1030A potentiostat (CH Instruments, Inc., Austin, TX, USA) through a custom-made high-speed electrical slip ring with operating speed range of 0-2500 rpm and dynamic resistance < 0.01 Ω (Model: SRS110073, Penlink, Sweden). The potentiostat was connected to the stationary part of the slip ring, whereas the moving part was coupled to the custom-made printed circuit board (PCB). The gold plated spring-loaded pins (Mill-Max Mfg. Corp., Oyster Bay, NY, USA) soldered to the PCB, formed the connection to the electrodes patterned on the disc (Figure 1B). The wires soldered to the PCB (Figure 1B, inset) enabled the connection to the potentiostat via the slip ring (Figure 1A).

Figure 1. (A) Photograph of a fully assembled experimental setup (slip ring connection to spring-loaded pins through PCB in the inset). (B) Exploded schematic view of the magnetic clamping system. (C) Photograph of the microfluidic disc and PCB with the magnetic clamping.
As can be seen in Figure 1B, the easy coupling of the PCB to the LoD device was achieved by incorporating rod magnets of Ø 4 mm and 3 mm height (Webercraft GmbH, Gottmadingen, Germany) in the poly(methyl methacrylate) (PMMA) layer (1 mm) glued to the PCB and in an additional PMMA layer (2 mm) placed below the LoD (Figure 1B). The slip ring was held in place by a 3D printed (Printer: MakerBot Industries, LLC, Brooklyn, NY, USA) holder made of polylactic acid (Reprap Aps, Copenhagen, Denmark), which was fixed on top of the disc mounted on the spin-stand using Ø 2-mm screws (Figure 1A and 1C).

2.4 Design, fabrication and assembly of microfluidic LoD device

The parts of the LoD device were designed using the computer aided design (CAD) software SolidWorks (Dassault Systèmes SolidWorks Corporation, Waltham, MA, USA). Based on the CAD files, the optical grade PMMA (Evonik Industries AG, Darmstadt, Germany) and double-sided pressure sensitive adhesive (PSA) tape (140 µm thick, ARcare® 90106 from Adhesive Research Ltd., Limerick, Ireland) were cut using a computer-controlled Epilog Mini 18 laser cutter equipped with a 30 W CO₂ laser (Epilog Laser, Golden, CO, USA) and Cameo 3 blade cutter (Silhouette, Inc., Lindon, UT, USA), respectively. The whole LoD assembly comprised four PMMA sheets, which were bonded to each other using a PSA, tape used interspersed between 4 layers of PMMA sheets (Figure 2A).

Figure 2. (A) Exploded view of the microfluidic disc with electrodes patterned on the bottom PMMA sheet. (B) Photograph of the microfluidic disc (details of a filtration and detection unit with electrode array in the inset). (C) Schematics of the cross section of the microfluidic layout (the arrows show the flow direction during rotation): sample (a) before and (b) after on-disc filtration.
The second PSA layer also allowed incorporation of the microporous cellulose acetate membranes (Tisch Scientific, North Bend, OH, USA) with 0.2 µm pore size. The filter membranes were custom cut using the blade cutter to fit the top of the collection chamber and placed between the PMMA sheets to separate the loading chambers from the collection chambers (Figure 2A, C). The turbid cell suspension was loaded into the loading chamber (Figure 2C(a)) and the clear cell-free suspension was driven to the detection chamber (Figure 2C(b)) after filtration achieved by rotating the LoD device at 700 rpm for 5 minutes. The PMMA sheet incorporated the eight gold electrode arrays (Figure 2B) each comprising of a working (WE), counter (CE), and pseudo-reference electrode (RE). The electrodes were patterned on the bottom PMMA substrate of the disc assembly using e-beam evaporation (20 nm Cr adhesion layer/200 nm Au) through a steel shadow mask, fabricated with the laser micromachining instrument as described previously (Amato et al., 2015).

2.5 Electrochemical measurements
All the electrochemical measurements were performed at room temperature with an eight-channel potentiostat (CHI1030A). For electrode characterization, cyclic voltammograms (CV) were recorded from -400 mV to 400 mV vs. Au pseudo-RE in 10 mM ferri/ferrocyanide using PBS (pH 7.4) as supporting electrolyte. Cleaning of the electrodes was carried out in 50 mM KOH dissolved in ultrapure water, with potential sweep from -200 mV to -1200 mV vs. Au pseudo-RE (scan rate 50 mV s⁻¹), performing at least three sweeps. pHCA detection was performed using square wave voltammetry (SWV) with a step potential of 0.001 V, square-wave frequency of 10 Hz and amplitude of 0.05 V. The applied potential ranges were 150-800 mV (control (J786) supernatant), 100-600 mV (M9 buffer), and 100-700 mV (M9 bacterial culture medium). The slip ring facilitates measurements during rotation as demonstrated by Andreasen et al. (Andreasen et al., 2015) yet, the electrochemical measurements presented in this paper were performed under static condition.

2.6 HPLC analysis
pHCA detection in 5 µL injection volumes was performed using a Waltham HPLC system (Thermo Scientific, MA, USA) equipped with a UV detector (333 nm) as previously described (Jendresen et al., 2015). The separation was achieved in a Discovery HS F5 column (3 µm particle size, 15 cm x 4.6 mm) at 30°C under gradient elution with a two–solvent system (A: 10 mM ammonium formate (pH 3.0 adjusted with formic acid) and B: acetonitrile) at a flow rate of 0.7 mL/min. The elution started at 5% B followed by linear increase of the fraction to 60% between 1.5 and 7 min after injection. The fraction of B was decreased back to 5% between 9.5 and 9.6 min, remaining there until 12 min.

2.7. Data analysis
For the calibration curves, the peak heights were obtained from the original data files using the CHI data analysis software and plotted against the corresponding concentration using OriginPro 2015 (OriginLab Corporation Northampton, MA, USA). The presented SWVs were background-subtracted
and baseline-corrected to improve the visualization and identification of peaks over the baseline without introducing any artifacts.

3. Results and Discussion

3.1 Electrochemical characterization and re-usability of the on-disc electrodes

Metal electrodes are most commonly fabricated on silicon or glass substrates although there are reports describing electrode fabrication on plastic substrates aiming at creating low-cost (Xu et al., 2011) and easy to integrate electrodes for microfluidic applications (Matteucci et al., 2016; Šnita et al., 2009). Patterning electrodes on plastic substrates can be achieved in a cost-efficient way by screen printing (Metters et al., 2011) or by using a stencil or shadow mask in combination with e-beam evaporation (Amato et al., 2015; Xu et al., 2011). However, it may be challenging to fabricate electrodes capable of providing a reproducible electrochemical signal as well as a stable metal (e.g. gold) layer, which is crucial for sensor development. The performance and reproducibility of the electrodes fabricated on PMMA substrate were evaluated by acquiring CVs. As shown in Figure 3A, the eight electrodes on the same disc have a peak potential separation ($\Delta E_p$) of 98 ± 2 mV and the ratio between the reduction and oxidation peaks was 1.07 ± 0.01. In addition, a linear relation between the oxidation and reduction peak currents and the square root of the scan rates was observed as presented in Figure 3B for sixteen electrodes from two discs. The recorded CVs indicate a quasi-reversible behavior of the redox couple on the electrodes and an excellent electrochemical reproducibility of the electrodes on the same disc as well as between discs. Although the observed $\Delta E_p$ was greater than 59 mV, which is the theoretically expected value for a reversible single-electron transfer reaction such that of ferri/ferrocyanide (A.J. Bard, 2001), the values are comparable with those reported for gold electrodes fabricated on glass (Andreasen et al., 2015), plastic (Matteucci et al., 2016) and silicon substrates (Zor et al., 2014).

![Figure 3](image-url)  
**Figure 3.** (A) CVs for 8 electrodes on the same disc acquired in 10 mM ferri/ferrocyanide using PBS (pH 7.4) as supporting electrolyte (scan rate 50mVs$^{-1}$). (B) Oxidation and reduction peak currents as a function of the square root of the scan rate (10-250 mVs$^{-1}$). The error bars represent the standard deviation of 16 electrodes among two discs (8 electrodes per disc). (C) Representative CVs for one-electrode acquired in 10 mM ferri/ferrocyanide before and after pHCA detection (scan rate 50mVs$^{-1}$).
Electrochemical oxidation of pHCA leads to the formation of a polymeric film on the electrode, impeding diffusion of the analyte and its oxidation on the electrode surface (Janeiro et al., 2007). In order to be able to re-use the electrodes after pHCA detection, cleaning and regeneration of the electrodes were evaluated. As shown in Figure 3C, the oxidation and reduction peaks of ferri/ferrocyanide are not clearly detectable after pHCA detection. There are various physical and chemical methods available for cleaning gold surfaces (Fischer et al., 2009). Due to the fact that the electrodes were fabricated on the plastic substrate, the optimal electrode cleaning process proposed by Fisher et al. (Fischer et al., 2009) proved to be unsuitable due to electrode delamination. However, the electrode surface could be cleaned by linear sweep voltammetry in 50 mM KOH solution in the potential range between -200 and -1200 mV (vs. on disc Au pseudo RE) at the scan rate of 50 mV/s as previously shown by Zor et al. (Zor et al., 2014). When CVs recorded for sixteen clean electrodes on two discs were compared with electrodes cleaned after pHCA detection we observed a slight increase in $\Delta E_p$ (from 98 ± 2 to 111 ± 9 mV) as well as decrease in the oxidation and reduction peak currents (from 421 ± 6 to 398 ± 11 µA and -392 ± 7 to -361 ± 16 µA, respectively). The increase in $\Delta E_p$ and decrease in the oxidation and reduction peak currents indicated that the electrochemical performance of the electrodes was not completely regained after pHCA detection. However, due to the good on-disc electrode reproducibility, the slight decrease in electrode performance can be overcome by performing on-disc calibration prior to determining the pHCA content in real samples.

3.2 pHCA quantification in supernatant

The electrochemical oxidation of pHCA was thoroughly investigated by Janeiro et al. using various electrochemical techniques (Janeiro et al., 2007). SWV was used in this work for quantification of pHCA because it is widely used in the development of electrochemical sensors, given its sensitivity and reliability (Chen and Shah, 2013). Detecting pHCA in cell culture supernatant could be a challenge due to the presence of possible interfering compounds that may be electrochemically active at the same potential as the target analyte. As a first step in the development of the detection method, pHCA was determined in M9 buffer (Figure S1A) and M9 culture medium (Figure S1B). However, our goal was to determine pHCA directly in bacterial supernatants, therefore we evaluated pHCA detection in a sample matrix that is very similar to the real samples, namely supernatant from the control (non pHCA producing) E. coli strain (CBJ786) collected after 24 h culture. The control strain has similar growth kinetics as CBJ792 and CBJ800 strains, but does not produce pHCA (Figure S2A). Signals recorded for 500 µM pHCA added to M9 buffer, M9 medium and to the bacterial culture supernatant of the control (CBJ786) are shown in Figure S1A, B, C. The observed difference in the pHCA oxidation potential is probably due to the pH dependent electrochemical oxidation of pHCA (Janeiro et al., 2007) since the pH of the supernatant at 24 h is close to pH 5.8, whereas the pH of M9 medium is 6.8, and of the M9 buffer is 7.4. We found that there are no electrochemical interferences in the SWV recorded between 150 and 800 mV in the range where pHCA is detected in control supernatant and therefore it was further used for the construction of calibration curves. Figure 4A shows representative SWVs for pHCA in the concentration range from 125 µM to 3 mM recorded in spiked control bacterial
supernatant. Figure 4B shows the linear part of the calibration curve. The pHCA concentrations in supernatants of recombinant microorganisms commonly reach 0.5 – 10 mM under laboratory conditions (Jendresen et al., 2015; Rodriguez et al., 2015; Xue et al., 2014). Therefore, the sensor array seems to be suitable for determining the analyte either directly or after dilution.

**Figure 4.** (A) SWVs of increasing pHCA concentration in control bacterial supernatant, potential step 0.001 V, frequency 10 Hz, amplitude 0.05 V. (B) Linear part of pHCA calibration curve recorded from spiked control bacterial supernatant, the error bars represent the standard deviation, n=4.

### 3.3 On-disc cell filtration and pHCA detection from real samples

Electrochemical sensors are prone to fouling (Hanssen et al., 2016). Therefore, the effect of the presence of bacterial cells on pHCA detection was investigated. In a typical bacterial culture, the cell density, determined as OD at 600 nm wavelength, can change from less than 0.2 to more than 10 (S2A). When determining pHCA in the presence of cell density, corresponding to OD600 of 2.0 after 9 h of growth, the recorded signal was not influenced by the presence of cells (S3). However, in a 24 h cell culture with high cell density, the electrochemical signal was 30 to 50% lower due to the sedimentation of cells on the electrode that caused electrode fouling (Figure 5). Since we aimed at determining the pHCA produced at the end point of the bacterial culture (24 h), a cell filtration unit (0.2 µm pore size) was integrated on the LoD platform as described in the Materials and methods section 2.4. The on-disc filtration was carried out after filling the loading chamber with 120 µl cell suspension followed by spinning the disc at 700 rpm for 5 minutes. The centrifugal forces push the filtered cell suspension to the detection chamber incorporated with the electrode at the bottom as it is shown in Figure 2C. The detection chamber was specifically designed by connecting it to a metering chamber as depicted in Figure 2B to enable the filling of the detection chamber with a defined volume which is independent of the filtration rate.

CBJ792 and CBJ800 *E. coli* strains, carrying derivative plasmids encoding TAL and the CBJ786 control strain were grown for 24 h in M9 medium supplemented with 2 mM Tyr. The pHCA production efficiencies of the genetically modified strains depend on their ability to convert Tyr into
pHCA as they express different recombinant TALs (Jendresen et al., 2015). After 24 h, almost the entire amount of Tyr was metabolized and or converted into pHCA (S2B, S2C). Therefore, Tyr did not interfere with the electrochemical detection of pHCA.

The pHCA content in the genetically modified E. coli cultures were measured at the end of bacterial growth at 24 h in bacterial culture supernatant. Cells were removed by on-disc filtration and the electrochemical detection was carried out under static condition after the detection chamber was filled. The pHCA concentration was calculated based on the peak current values of the recorded SWVs using the calibration curve. The electrochemically determined pHCA content in centrifuged and on-disc filtered samples was found to correlate well with the data obtained from HPLC analysis (Figure 5).

4. Conclusion

Monitoring of biological processes is important when developing new genetically modified strains or evaluating the quality of an already well-known process. In this work, we used pHCA producing E. coli as a model system for developing a LoD device for bioprocess monitoring. Due to the lack of natural availability of and the demand for plant secondary metabolites in the manufacturing of healthcare and nutritional products, microbial organisms are often engineered to contain artificial biosynthetic pathways. We presented a robust, easy to use and compact electrochemical LoD device with integrated filtration for detection of pHCA produced by genetically modified E. coli strains as an alternative approach to traditionally used methods for pHCA quantification. The gold electrodes fabricated on plastic (PMMA) substrate proved to be stable and reusable as well as facilitates reproducible measurements. The electrochemical quantification of the pHCA content at the end of the bacterial culture measured for two E. coli strains was in a good agreement with values obtained from HPLC.
analysis. The presented device can be further developed for time-dependent monitoring of pHCA content by including additional sample pre-treatment steps to enable pHCA detection in the presence of high Tyr concentration. In addition, by further parallelization to facilitate higher detection throughput, the sensing platform has a great potential for being implemented as a compact analytical tool for bioprocess monitoring.

Acknowledgements

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References


Figure captions:

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**Figure 5.** pHCA quantification in presence of cells, on-disc filtered and centrifuged supernatant vs. HPLC, the error bar represents the standard deviation between measurements, n=3.
Supplementary material

Lab-on-a-disc platform for screening of genetically modified *E. coli* cells via cell-free electrochemical detection of *p*-Coumaric acid

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S1. SWVs for 500 µM pHCA in M9 buffer (a), M9 culture medium (b) and CBJ786 control supernatant (c) with respective blanks.

S2. Changes in three different *E. coli* stains during 24h of culture: Growth (a); Tyr consumption (b) and pHCA production (c).
S3. Comparison between signal recorded for 500 µM pHCA in the presence (OD=2) and absence of cells.